

Computer Algebra and Differential Equations An Overview

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Abstract

We present an informal overview of a number of approaches to differential equations which are popular in computer algebra. This includes symmetry and completion theory, local analysis, differential ideal and Galois theory, dynamical systems and numerical analysis. A large bibliography is provided.

1 Introduction

Differential equations represent one of the largest fields within mathematics. Besides being an interesting subject of their own right one can hardly overestimate their importance for applications. They appear in natural and engineering sciences and increasingly often in economics and social sciences. Whenever a continuous process is modeled mathematically, chances are high that differential equations are used.

Thus it is not surprising that differential equations also play an important role in computer algebra and most general purpose computer algebra systems provide some kind of `solve` command. Many casual users believe that designing and improving such procedures is a central problem in computer algebra. But the real situation is somewhat different. Many computer algebra applications to differential equations work indirectly; they help to study and understand properties of the solution space.

The purpose of this article is to sketch in an informal way some of the main research directions in this field and to provide a starting point for more detailed studies by giving a large number of references. Therefore we omit all mathematical details (there is not a single formula in this article!) but describe briefly the central ideas. For the same reason we often cite introductory articles or books and not the historically first or the most “ground breaking” work.

The bibliography is of course far from being exhaustive. As a further source of references one should mention the survey by Singer [115]. It gives much more details, especially on the more algebraic approaches, and contains a large bibliography. The same holds for the more focused surveys by Hereman [57, 58] covering symmetry theory and related fields and the one by MacCallum [74] on the integration of ordinary differential equations. In addition there have been three conferences devoted exclusively to differential equations and computer algebra. Their proceedings [114, 123, 124] contain a number of useful introductory or review articles on more specialized topics.

We are taking a rather broad view and consider more or less any constructive approach to differential equations as “computer algebra”. This also implies that we do not pay special attention to implementations. Among the many different approaches to differential equations which fall under this broad definition of computer algebra one can distinguish certain directions which have found most attention (at least measured in the number of articles devoted to them). We concentrate in this article on the following eight topics: (i) *solving* differential equations, (ii) *local analysis*, (iii) *symmetry analysis*, (iv) *completion*, (v) *differential ideal theory*, (vi) *differential Galois theory*, (vii) *dynamical systems theory*, and (viii) the relation between *numerical analysis* and computer algebra.

A comparison of the impact made by symmetry analysis and by differential Galois theory, respectively, demonstrates the importance of computer algebra tools. The latter one is a hardly known theory studied by a few pure mathematicians. The former one remained in the same state for many decades following Lie’s original work. One reason was definitely the tedious determination of the symmetry algebra. As soon as computer algebra systems emerged, the first packages to set up at least the determining equations were written. Since then Lie methods belong to the standard tools for treating differential equations.

2 Solving Differential Equations

Most computer algebra systems can solve some differential equations. They mainly apply standard techniques like those in Zwillinger’s handbook [138] or try “pattern matching” in a list of solved equations like Kamke [64]. Heuristics often extend the applicability of this approach, for example by finding a transformation of a given equation to one that can be handled by the implemented methods.

Although this approach solves more differential equations than one might expect (see e.g. the recent review by Postel and Zimmermann [98]¹), it has some drawbacks. A major one is that no information is obtained, if the computer algebra system does *not* return a solution. It could be that the

¹An updated version can be found at <http://www.mupad.de/BIB/poszim96.html>.

given differential equation has indeed no solution (in closed form) or that simply the heuristics could not determine a suitable transformation.

For that reason researchers in computer algebra are especially interested in *decision algorithms*. These either yield a solution in a specific class of functions or decide that no such solution exists. However, so far only for linear ordinary differential equations such algorithms are known. There it is possible to decide with the help of differential Galois theory (see Sect. 7) whether or not Liouvillian solutions exist.

There exists a number of reasons for this perhaps disappointing situation. Computability theory yields principal limits to what can be solved. For example if one restricts to *computable functions* some classical existence theorems for differential equations fail [1, 99]. More precisely, one has constructed examples of differential equations where one can show that solutions exist but that it is not possible to compute them. Some further (positive and negative) results in this direction can be found in [29].

Ideally, a solution algorithm should return the *general solution*. But for nonlinear equations it is surprisingly difficult even just to define this term. A rigorous resolution of this problem (for ordinary differential equations) based on differential ideal theory (see Sect. 6) was only recently presented [60].

Intuitively one would expect that the general solution depends on some arbitrary parameters (constants or functions) and that every solution of the differential equation can be obtained by a suitable specialization of these. This works fine for linear equations where the solution space has the structure of a vector space. But many nonlinear equations possess in addition *singular integrals* not contained in the general solution. A solution algorithm should probably automatically compute these, too.

Similarly, defining the term “closed form solution” is notoriously difficult. Is a solution in terms of, say, Bessel functions in closed form or not? Up to now no generally accepted definition has emerged. The basic idea behind “closed form” is that of *finite constructibility* out of a set of “elementary functions”; but now the problem arises how to define “elementary”. Note that this is an algebraic and not an analytic property!

On the practical side one must see that even if a solution in closed form can be computed it may take very long and the result may be completely useless, as it is too large. Especially, if the main goal is to obtain an impression of the behavior of the solution, it is often much more efficient to resort to numerical methods. For that reason many computer algebra systems provide at least for ordinary differential equations some standard numerical integrators like Runge-Kutta methods etc. (see also Sect. 9).

In any case one can state that a notable solution theory exists only for ordinary differential equations (see e.g. the survey [74]), mainly based on either Lie symmetry theory or differential Galois theory. But the former one often does not yield complete algorithms, so that one must resort to heuristics in intermediate steps. The algorithms of the latter one suffer from

a very high complexity and are in practice often rather useless, especially for higher order equations. We will discuss these problems in a bit more detail in Sects. 4 and 7.

Another possibility that also addresses the problem of useless output is to search only for “simple” solutions [9]. Popular variants are polynomial, rational [3] or exponential [14, 93] solutions. Because of their simple structure it is often possible to determine such solutions, if they exist, rather fast. But one should note that the classical methods for their computation are not always useful for computer algebra. It is still an active field of research to design effective algorithms being able to handle larger examples.

For partial differential equations the situation is much worse; usually one must already be happy, if one can find any closed form solution at all. In the last century mathematicians designed some solution methods (see e.g. the survey [129]). However, most of them are more or less forgotten; at least they are no longer found in standard textbooks. It could be quite interesting to revive some of them for use in computer algebra systems.

There exist a few implementations of standard techniques like characteristics, separation of variables or integral transforms (see for example [23]), but they can usually handle only rather simple equations. Often they just reduce the partial differential equation to a system of (nonlinear) ordinary differential equations and the question is whether this can be solved. The most important approach to constructing solutions of partial differential equations is provided by symmetry theory (see Sect. 4).

One can argue whether it really makes sense to speak of the general solution of (a system of) partial differential equations. For example one definition of a harmonic function is that it solves the Laplace equation (or more generally all holomorphic functions are solutions of the Cauchy-Riemann equations). Thus one might prefer to say that the Laplace equation defines a class of functions.

In some simple cases like the wave equation one can give an explicit parameterization of this class in terms of some arbitrary functions which one may call the general solution. But usually no such parameterization exists. In order to get a well-defined problem one must prescribe some initial or boundary conditions. In most applications such conditions arise automatically anyway.

3 Local Analysis

If it is not possible to construct a closed form solution, one may go for an approximate solution describing the behavior of the solution in the neighborhood of a given point.² At *ordinary points* a Taylor series suffices; at

²Especially in the linear case local solutions can also be very useful for the construction of closed-form solutions.

singular points more general expansions must be used. Local analysis is essentially a complex theory, even if one studies only real equations. Especially, if one wants to determine the radius of convergence of a formal solution, one must also take the complex singularities into account.

In the case of linear differential equations singular points are only possible at singularities of the coefficients. Therefore one speaks of *fixed* singularities. Using the *Newton polygon* of the associated differential operator they can be further classified into *regular* and *irregular* ones [31].

In the neighborhood of regular singular point one can represent the solution in form of a *Frobenius series*, a polynomial in $\log x$ with Taylor series coefficients multiplied by a factor $(x - x_0)^\alpha$ with α a complex number. At irregular singular points the solution has typically an essential singularity. It varies so rapidly that it makes no sense to construct an approximation; instead one tries to capture its asymptotic behavior which requires the addition of an exponential part. An elementary introduction can be found in the textbook [10].

There exist various algorithms for the construction of approximate or asymptotic solutions, partly dating back at least to Frobenius. Some are discussed together with implementations in [94, 126]. A main problem in their application is that one cannot use approximations of the location of the singularities. One must not only solve polynomial equations but in general work with algebraic numbers which is quite expensive in any computer algebra system. But a careful analysis of the algorithms can often significantly reduce the necessary amount of computations with algebraic numbers.

Recent work concerns an extension of the theory to first order systems [8, 93]. In principle, one can transform any system into a single equation of higher order, e. g. using *cyclic vectors*. But this approach is rather inefficient, especially in higher dimensions. Hence one is interested in dealing directly with systems. *Moser's algorithm* performs here the classification into regular and irregular singularities; a rational version of it avoiding the use of algebraic extensions was presented by Barkatou [7].

For nonlinear differential equations the situation becomes much more complicated as *spontaneous* or *movable* singularities may occur, i. e. their location depends on initial or boundary data. One usually speaks of the *Painlevé theory* [26, 61, 69]. It was introduced by Painlevé while searching for new special functions and there still exists a strong connection to special function theory. If all singularities are poles, no branch points appear in the (general) solution and it is single valued. A differential equation without movable branch points is said to possess the *Painlevé property* or to be integrable in the sense of Painlevé.

In general, it is not possible to check algorithmically whether or not a given differential equation has the Painlevé property. But there exist methods to check at least some necessary conditions; such methods are usually called *Painlevé test*. The main approach consists in trying to construct a

Laurent series around the singularity. Essentially, the test is passed, if this expansion has sufficiently many *resonances* or *Fuchsian indices* (free coefficients) to represent the general solution and if these occur at non-negative powers. In the case of negative resonances a perturbation approach [27] yields further information. Some references concerning implementations can be found in [106].

Weiss *et al.* [133] generalized the Painlevé theory to partial differential equations where a whole singularity manifold must be considered. This extension is much used in the theory of integrable systems, as the Painlevé test represents an important indicator for (complete) integrability and can be performed comparatively easily. The *Painlevé conjecture* states that every ordinary differential equation obtained by symmetry reduction (see Sect. 4) of an integrable system is of Painlevé type; only weakened versions of it have been proven [2, 80]. Truncated series expansions are useful for constructing Bäcklund transformations, Lax pairs and much more [132]. There also exist relations to non-classical symmetry reductions [36].

Comparing with the linear case we see that the nonlinear theory described so far corresponds roughly to the case of a regular singular point, namely a Frobenius expansion of the general solution (often restricted to integer α). Extensions to irregular singular points have been proposed for the nonlinear case by Kruskal *et al.* [69]. Essentially the strategy is the same: one adds an exponential part to the solution ansatz.

4 Symmetry Analysis

Of all the approaches discussed in this overview, symmetry analysis has made the strongest impact on computer algebra applications to differential equations. The most general definition of a symmetry is that of a transformation that maps solutions into solutions. Depending on the kind of transformations considered one obtains different kinds of symmetries. One possible application of symmetries is the construction of (special) solutions. Other goals are classifications, a proof of (complete) integrability, separation ansätze, conservation laws and much more. Several excellent textbooks on this subject are available, e.g. [12, 86, 122].

Symmetry analysis goes back to the seminal work of Lie. He developed the concept of Lie groups in his quest for a Galois theory for differential equations. As we will see later in Sect. 7, not much has remained of this original motivation. Symmetry and Galois theory have developed in very different directions. Even the relation between the Lie symmetry group and the Galois group of a linear differential equation is rather unclear.

The most popular variant of symmetry analysis deals with *Lie point symmetries*. They are generated by vector fields acting on the space of independent and dependent variables. These vector fields span the Lie algebra

of the Lie group of symmetries. The decisive observation of Lie was that for most purposes it suffices to work with the vector fields (or infinitesimal symmetries) instead of the symmetries themselves. This leads effectively to a linearization of the problem.

The symmetry generators arise as the solutions of a linear system of partial differential equations, the so-called *determining system*. For ordinary differential equations it is unfortunately sometimes as difficult to solve this system as to solve the original one. This holds especially for first order equations where the original equation is just the characteristic equation of the determining equation. For partial differential equations the determining system is typically very over-determined and contains often some trivial equations allowing in many cases a rather straightforward solution. The analysis of determining systems is a typical task for completion algorithms (see Sect. 5).

For ordinary differential equations the existence of a sufficiently large and *solvable* symmetry algebra implies that its general solution can be constructed by quadratures, as each symmetry allows us to reduce the order of the equation by one (“cascade integration”). In the case of partial differential equations symmetry reductions lead only to special solutions, namely those being invariant under the symmetry group. Here each symmetry allows us to transform to an equation with one independent variable less. Thus with a sufficiently large solvable symmetry algebra a partial differential equation can be reduced to an ordinary differential equation.

At intermediate steps of the reduction linear partial differential equations must be solved. This starts with the determining system. Later, in order to obtain the reduction, one must either perform a coordinate transformation such that the symmetry generator is rectified (*canonical coordinates*) or the *differential invariants* of the symmetry must be determined. These are functions annihilated by the symmetry generator, i.e. they are defined as the solutions of a linear partial differential equation.

Thus the usefulness of Lie symmetries depends crucially on the ability to effectively solve all the arising partial differential equations. At first sight it might look, as if, especially for ordinary differential equations, we made the problem only worse. But in many cases of practical interest it turns out that is much easier to solve these linear partial differential equations than the original equation. Often the repeated application of rather simple heuristics suffices to completely solve at least the determining system.

There exist so many implementations of symmetry methods that it is rather difficult to keep an overview; we refer again to the surveys by Herman [57, 58] with their huge bibliography. In almost any computer algebra system one can find at least a package for setting up the determining system which sometimes also tries to solve it. Some packages are even able to use the symmetries to compute automatically closed form solutions for some classes of differential equations.

Although Lie point symmetries proved to be very useful in many applications, there still exist many differential equations of practical interest which do not possess symmetries or at least not sufficiently many. There are two basic approaches to generalize the theory. One can consider more general transformations; this leads to *generalized* or *Lie-Bäcklund symmetries* [5]. Alternatively, one weakens the requirement that every solution is mapped into a solution; this yields the so-called *non-classical methods*.

The generators of generalized symmetries may also depend on derivatives of the dependent variables. The corresponding symmetry transformations are now no longer given as a simple exponential flow but must be determined as solutions of a partial differential equation. In principle, the determination of generalized symmetry proceeds exactly as for point symmetries. However, a bound on the order of the derivatives appearing in the generator must be chosen at the beginning of the computation. Thus it is not possible to algorithmically construct all generalized symmetries with this approach.

Generalized symmetries are of much interest for (completely) integrable systems [38, 134]. The existence of a *recursion operator* or a *master symmetry* generating an infinite hierarchy of symmetries is a strong indication that the considered system is integrable. This approach also circumvents the problem of the a priori bound for the order of the generator. Reduction with respect to generalized symmetries is an important tool for the construction of *soliton solutions*. It is also possible to classify nonlinear partial differential equations using these symmetries [82].

Non-classical reductions can be understood within the general scheme of augmenting a given differential equation with *differential constraints* [87]. This corresponds to requiring that only some solutions are mapped into solutions. Hence one hopes to find more symmetries (these are sometimes called *weak* or *conditional symmetries*). In this approach the emphasis lies less on group theory but on the theory of over-determined systems of partial differential equations and thus on questions of completion (cf. [111]).

The first non-classical method was developed by Bluman and Cole [11] and uses the invariant surface condition as constraint. Although this leads for many differential equations to new reductions, the drawback is that the determining system becomes nonlinear. The *direct method* of Clarkson and Kruskal [25] tries to reduce a given partial differential equation to a system of ordinary differential equations by constructing a good ansatz; it corresponds to a special case of the method of Bluman and Cole.

The main problem in the method of differential constraints is to find compatible constraints leading to non-trivial reductions. Besides using the invariant surface condition no systematic way has been discovered so far and thus it remains essentially a game of “try and error”. For this reason differential constraints have not yet found much attention in applications.

One can also combine both generalizations and obtains then so-called “non-classical or conditional generalized symmetries”. These have been used

to study the interaction of traveling waves solution of non-integrable systems [39]. However, one must note that this approach does not lead beyond the theory of differential constraints. As already pointed out by Olver and Rosenau [89] any differential constraint may be considered as the characteristic of a generalized symmetry generator.

5 Completion

Most textbooks on differential equations treat only *normal* systems (or systems in *Cauchy-Kowalevsky form*). For ordinary differential equations this implies that one assumes that the equations can be solved for the highest order derivatives. For partial differential equations one must furthermore assume the existence of a distinguished independent variable such that one can solve for its derivatives to obtain the Cauchy-Kowalevsky form. However, in many fields one encounters systems of differential equations which are not normal. A simple example are the determining systems in symmetry analysis (see Sect. 4) which are usually over-determined. Non-normal systems also occur naturally in differential geometry and in theoretical physics (gauge theories).

For a non-normal system it is a priori not clear whether it possesses any solutions at all. It may happen that the system is inconsistent. This can only be decided after the construction of all *integrability conditions*. These are further differential equations satisfied by any solution of the system but nevertheless *algebraically* independent of it. While it is easy to construct one integrability condition (typically this requires only a cross-derivative), it is not so easy to decide when *all* have been found, as in principle an infinite number of conditions must be checked.

The process of finding all integrability conditions is called *completion*. It results in a *formally integrable system*,³ as after completion it is straightforward to construct order by order a formal power series solution. Under additional assumptions it is sometimes possible to show the convergence of the series. This leads for analytic systems to existence and uniqueness theorems like the Cartan-Kähler theorem (the well-known Cauchy-Kowalevsky theorem is a special case of it). For non-analytic systems solvability is a much more complicated question due to Lewy type effects [72].

The first systematic approach to the problem of completion was probably provided by the Janet-Riquier theory [62] with the introduction of *passive* systems. Their definition is based on a *ranking* of the derivatives which decides in what order the integrability conditions are constructed. The completion can be done completely automatically only for quasi-linear systems (if all arising integrability conditions are also quasi-linear), as it must be

³Not to be confused with a *completely* integrable system as mentioned in Sect. 4!

possible to solve for the leading derivative. In this case the resulting passive system is sometimes called a *standard form* [103].

In geometric theories the notion of a passive system is replaced by *involution*. It combines a geometric definition of formal integrability with an algebraic criterion for the termination of the completion. As an intrinsic concept involution requires no coordinate dependent ingredients like a ranking. Involution analysis based on the Cartan-Kähler theory [17] for exterior systems is discussed from an algorithmic point of view in [55, 56]. A completion algorithm for the jet bundle formalism based on the formal theory of Pommaret [96] was presented in [108].

Completion algorithms are very useful in the symmetry analysis of differential equations. Once a system is either passive or involutive, one can make statements about the size of the solution space [103, 109]. Thus it is possible to compute the size of the symmetry group without explicitly solving the determining system or to determine the loss of generality in a symmetry reduction [110]. One can even compute the abstract structure of the symmetry algebra without solving the determining system [73, 104].

These concepts are closely related to Gröbner bases in commutative algebra. This holds especially for the Janet-Riquier theory where rankings play a similar role as in the definition of a Gröbner basis. Therefore one sometimes finds the term *differential Gröbner basis* for a passive system. Integrability conditions arising from cross-derivatives may be considered as “differential S-polynomials”. But these analogies acquire a precise meaning only in the context of differential algebra (see Sect. 6).

There is a one-to-one correspondence between linear systems of partial differential equations in one dependent variable with constant coefficients and polynomial ideals. This has lead in commutative algebra to the new concept of an *involution basis* of an ideal [50]. It is computed using algorithms coming from the completion theory of differential equations, but it is an ordinary (though not reduced) Gröbner basis. In some cases the new algorithms are considerably faster than the classical Buchberger algorithm. Involution bases also allow for a straightforward determination of the *Hilbert polynomial*⁴ [6].

6 Differential Ideal Theory

Differential ideal theory belongs to the field of *differential algebra*. It can be informally described as an attempt “to write differential in front of everything in algebra”. It deals with differential rings, differential fields etc. This requires an algebraic definition of differentiation. In differential algebra any mapping that is linear with respect to addition and satisfies the Leibniz or

⁴For a discussion of the Hilbert or *differential dimension polynomial* of differential equations see [90, 109].

product rule is called a *derivation*. A differential ring is a commutative ring together with one (or more) derivation.

Differential polynomials arise by adjunction of differential indeterminates to a differential ring. However, adjoining one differential indeterminate corresponds to adjoining infinitely many algebraic indeterminates, as one must introduce all its derivatives as additional, algebraically independent variables. Thus Hilbert's basis theorem does not apply and the ring of differential polynomials is not Noetherian.

A *differential ideal* is an ideal which is in addition closed under the derivation of the differential ring. Many of the basic ideas in differential ideal theory can be traced back to Ritt [105]; the most advanced book is still the one by Kolchin [68]. The *Ritt-Raudenbush theorem* asserts that any perfect differential ideal, i.e. one that is equal to its radical, is the radical of a finitely generated differential ideal. In analogy to algebraic geometry one can try to introduce *differential algebraic varieties* as the set of "zeros" of a system of differential polynomials, i.e. the solution set of the corresponding differential equations. In general, this requires an extension of the base field.

As in the purely algebraic theory one would like to introduce Gröbner bases. But as the Ritt-Raudenbush theorem is weaker than full Noetherianity, algorithms along the lines of the Buchberger algorithm do not always terminate [21]. More generally, one can prove that the *ideal membership problem* is undecidable for arbitrary differential ideals [43]. However, this result is more of theoretical interest, as for finitely generated ideals the decidability is still an open question. In any case one must say that no generally accepted definition of a differential Gröbner basis has yet emerged.

There exist two basic strategies to circumvent this principal problem. One can either restrict to special ideals where a proof of termination is possible or one weakens the properties expected of a differential Gröbner basis. The completion algorithm of the Janet-Riquier theory (see Sect. 5) can be considered as a simple example for the first strategy. An example for the second one are the bases introduced by Mansfield [79]. They are computed with pseudo-reductions and have thus weaker properties than their algebraic counterpart. Especially, it may happen that one leaves the ideal.

Recently, Boulier *et al.* [13] presented a *Rosenfeld-Gröbner algorithm* which computes a representation for perfect differential ideals in the following form. The ideal is written as a finite intersection of saturations ideals; these are radical differential ideals defined by a system of differential polynomial equations and inequalities. This representation allows for an easy algorithmic test of radical ideal membership and for computing formal power series solutions.

Open problems are to obtain a minimal decomposition, i.e. to use only a minimal number of saturation ideals, and to find bases for these ideals (avoiding the inequalities). These questions are closely related to the inclusion problem for differential ideals which in turn can be seen as the problem

of determining the relation between the singular and the general solutions of a differential equation [60]. The principal obstacle in the construction of the bases is a very typical one in differential algebra. A theorem of Ritt asserts that by taking sufficiently many derivatives of the equations one can always get a basis but no bound for the number of derivatives needed is known.

Differential algebra is applied in *automatic theorem proving* in differential geometry [137]. This is similar to the use of algebraic ideal theory in theorem proving in elementary geometry. For this kind of applications *characteristic sets* seem to be more useful than Gröbner bases. A nice example for the possibilities here is the automatic derivation of Newton's law of gravity from the three Kepler laws [136].

Besides ideals of differential polynomials there has been some work on ideals of linear differential operators or ideals of the Weyl algebra [42]. But here one is dealing with non-commutative rings. One could also consider the Cartan-Kähler theory (see Sect. 5) as a kind of differential ideal theory, as it represents differential equations by closed ideals of differential forms.

7 Differential Galois Theory

Already Lie was looking for a differential analogue of the (algebraic) Galois theory, when he introduced Lie groups. What is nowadays usually called *differential Galois theory* [75, 116] has however no connection to Lie symmetry theory. The latter one uses continuous transformation groups and can be applied to any differential equations. But as discussed in Sect. 4 it is not completely algorithmic. The former one is based on linear algebraic groups. It considers exclusively linear ordinary differential equations and culminates in various algorithms for explicitly computing Liouvillian solutions.

Determining the solutions of linear differential equations is a very classical topic and many famous mathematicians like Liouville, Fuchs, Klein or Jordan studied it in the last century and their results are still very important for the design of algorithms. Differential Galois theory was essentially founded by Picard and Vessiot and given its modern form by Kolchin [68]. Pommaret [97] developed an alternative theory following more closely Lie's ideas and using the formal theory.

We mentioned already in the last section that the solutions of algebraic differential equations typically lie in some extension of the base differential field. In differential Galois theory these extensions are studied in more detail. One can show that for a linear equation of order q a differential splitting field, the *Picard-Vessiot extension*, exists containing q solutions linearly independent over the constants. The *differential Galois group* consists of field automorphisms of this extension that commute with the derivation and that leave elements of the base field invariant.

Very important extensions of the field of rational functions are the *Liou-*

Liouvillian functions. They comprise essentially all expressions one can “easily write down”. Allowed operations are the usual arithmetic ones, roots, exponentials, logarithms, integrals and algebraic functions. A more formal definition uses a tower of simple extensions. An important point is that for any Liouvillian function one needs only a finite number of extensions, thus it is algorithmically constructible. Most expressions one would call “closed-form” are in fact Liouvillian.

Most solution algorithms are based on the seminal work of Singer [113]. He showed that the logarithmic derivative of any Liouvillian solution is algebraic and determined an a priori bound for the degree of the minimal polynomial, namely the Jordan bound for the index of an Abelian normal subgroup of a finite linear group. In principle, this suffices to determine all Liouvillian solutions, but the bound grows rapidly with the order of the equation leading thus to a very high complexity of the algorithm.

Using the representation theory of finite groups Ulmer [125] could significantly improve the bound given by Singer, so that at least the treatment of equations up to third order seems feasible, but there does not yet exist an implementation. Group theory yields also a number of other interesting results like criteria for the existence (and number) of algebraic solutions (the solutions which are most expensive to determine belong to this class) and gives the basic case distinctions in the solution algorithms.

The original work of Singer covered only equations with rational coefficients. Later, it was extended to Liouvillian coefficients [15, 117]. For second order equations Kovačič [63, 34] developed independently a solution algorithm. Only much later one could show that the classification behind this algorithm can also be derived within the Singer theory [119]. The Kovačič algorithm has been implemented in several computer algebra systems.

An alternative approach based on the *invariant ring* of the differential Galois group was presented by Fakler [37] following ideas going back to Fuchs (see also the work of Singer and Ulmer [119] and van Hoeij and Weil [127], respectively). For second order equations there exists an isomorphism between the invariant ring and the rational solutions of some symmetric power of the differential equation. This isomorphism allows one to derive explicit solution formulae and thus a rather efficient algorithm.

Determining the differential Galois group of a given equation is difficult. Some progress has recently been made for second and third order equations [118] where the problem could be reduced to finding solutions of some associated linear differential equations in the coefficient field and to factoring such equations. If there was an easy way to compute the group directly, one could probably design more efficient solution algorithms. But currently it is the other way round: the solution algorithms help finding the group.

There has also been some work on the *inverse problem* of differential Galois theory. Here a linear algebraic group is given and the task is to determine a differential equation that has it as differential Galois group. One

can prove that such a differential equation always exists [83]. Ramis [101] showed that it is often possible to reduce the inverse problem to the direct problem and then even give an explicit solution.

All the theory mentioned here works only for *irreducible* equations. Thus the (efficient) *factorization* of linear differential operators is an important problem in differential Galois theory. A solution of this problem based on the Newton polygon was recently presented by van Hoeij [126]. Factorization (although only of polynomials) is an issue in differential ideal theory, too.

Differential Galois theory can also be used to find (Liouvillian) *first integrals* [78, 100, 131]. These help to construct explicit solutions [77]. Other applications appear in the theory of (completely) integrable systems. Ziglin has given an algebraic characterization of such systems based on their monodromy group. His criterion for integrability can be rephrased in terms of certain properties of the differential Galois group [24, 84].

8 Dynamical Systems

Applications in dynamical systems theory are not really in the main stream of computer algebra. Conversely, numerical computations play a much more prominent role within dynamical systems theory than symbolic ones. Nevertheless, the use of computer algebra systems is becoming more and more popular in this field. Their main task is the determination of approximations or more generally perturbation analysis [102]. Two fundamental techniques in dynamical systems theory are especially well suited for computer algebra: normal forms and center manifolds. Other applications include bifurcation analysis, the Poincaré map and Hilbert's 16th problem.

Defining *normal forms* and deriving algorithms to compute them is a classical topic in computer algebra. For dynamical systems normal forms have already been introduced by Poincaré, Birkhoff, Gustavson and many others, often in the context of celestial mechanics [16, 30]. They form the basis for the solution of many problems in dynamical systems theory like for example stability or bifurcation analysis. One should however note that the word “normal form” is used here in a slightly different meaning than usually in computer algebra, as the normal form of a dynamical system is only an approximation of it.

The main idea behind normal forms is to study the system in the neighborhood of a *fixed point* (or *equilibrium*) and to try to remove by a near-identity coordinate transformation as many nonlinear terms from the defining vector field as possible. According to the Hartman-Grobman theorem all such terms can be eliminated near a *hyperbolic fixed point* where the Jacobian has no zero or purely imaginary eigenvalues [92]. Thus at such a point linear stability theory is sufficient.

Around other types of equilibria like centers the analysis is more in-

volved. This concerns especially Hamiltonian systems where fixed points can never be hyperbolic. If there are *resonances* between the eigenvalues of the Jacobian, the normal form is necessarily non-linear. In order to determine a normal form one makes a power series ansatz for the coordinate transformation and determines the coefficients of the ansatz by requiring that besides the resonances all non-linear terms of the vector field up to a certain order disappear. The resulting transformed field is a normal form.

Further complications arise, if the linear part of the vector field is not semi-simple, i.e. if it contains a nilpotent part. In this case an additional normalization is necessary, as the classical algorithms consider only the semi-simple part. This requires further tools from invariant and representation theory of Lie algebras. For the special case of $sl(2, \mathbb{R})$ this is discussed in some detail in [28]. For planar systems with nilpotent linear part generic normal forms up to eighth order have been computed in [40] using REDUCE.

An algorithm for computing normal forms that is suitable for implementation in a computer algebra system was presented by Walcher [130]. It is closely related to *Lie transforms* [81]. This technique has its origin in Hamiltonian mechanics where it yields a canonical transformation. However, it can be extended to general dynamical systems. In contrast to this Birkhoff normal form Gatermann and Lauterbach [48] took normal forms from singularity theory in order to study bifurcation phenomena. For equivariant systems (see below) they automatically classify them using Gröbner bases.

Computer algebra is also much used to determine (approximations of) *center manifolds* [20], a special form of invariant manifolds. If a dynamical system possesses a center manifold, it often suffices to study its behavior on this manifold. If the zero solution of the reduced system is stable, solutions of the original system for initial data sufficiently close to the center manifold will approach this manifold exponentially fast. Thus the reduced system completely describes the asymptotic behavior of such solutions.

Center manifold theory has such a great importance, because it yields a reduction of the dimension and thus often a considerable simplification of the analysis. Sometimes it is even possible to reduce an infinite-dimensional problem to a finite-dimensional one. There are two main computational steps. First we need an approximation for the center manifold, then we must compute the reduced system. As in normal form theory, this is done with a power series ansatz [41]. Laure and Demay [70] showed for the Couette-Taylor problem how computer algebra and numerical analysis can interact to solve a complicated bifurcation problem for an infinite-dimensional problem using a reduction to a finite-dimensional center manifold.

But also some classical (computer) algebraic problems are of great importance in the study of dynamical systems. For example, before a fixed point can be analyzed it must be determined. This requires the solution of a nonlinear system of algebraic equations. If the defining vector field is rational, this can be done with Gröbner bases. Often the vector field depends

on some parameters. At certain values of these parameters, the properties of the vector field may change, i. e. a *bifurcation* occurs. The determination of these values is a fundamental problem in dynamical systems theory.

Of special interest are here *equivariant systems* where the vector field is invariant under the action of a symmetry group [52, 53]. Here one can use linear representation theory and polynomial invariant theory for determining the fundamental invariants and equivariants [46, 135]. Using normal forms they enable the local bifurcation analysis, i. e. the typical bifurcation diagram in the neighborhood of a critical point can be derived.

If bifurcations of periodic solutions are to be studied, the *Poincaré map* is often a very useful tool. However, in general it is not possible to obtain it analytically. Thus one computes again a power series approximation of it. The bifurcation depends then on the Taylor coefficients. They satisfy a system of differential equations which must be integrated numerically. In order to set up this system one needs higher order derivatives of the vector field which can often be determined only by computer algebra. A combined numerical-symbolical approach to the Poincaré map is described in [66]. A pure MAPLE package using the built-in numerical and graphical facilities for plotting Poincaré sections was presented in [22].

A more theoretical application concerns *Hilbert's 16th problem* of bounding the number of limit cycles in a planar polynomial system. For quadratic systems a lot of results are known [107]; however already the cubic case becomes very complicated. An important subproblem is the *center problem*, namely to distinguish between a focus and a center. The derivation of sufficient and especially of necessary conditions for a center can be very involved and is sometimes hardly feasible without computer algebra [91]. In a recent study of cubic systems [35] a CRAY-J90 had to be used.

9 Numerical Analysis

It was already mentioned in Sect. 2 that the capabilities of computer algebra systems in explicitly solving differential equations are limited. This holds especially for partial differential equations. Hence numerical methods have lost nothing of their importance. Symbolic and numerical computations can interact in many ways and most systems provide some numerical facilities.

The oldest and simplest approach consists of *interfacing* a computer algebra system and a numerical library. Typically the interaction is one-way: the computer algebra system is used to derive the differential equations; the interface generates code in the language of the numerical library (perhaps including some optimization steps); finally, the differential equations are solved by some routines from the numerical library.

To some extend most common computer algebra systems can do this, as they provide commands to convert an expression into C or FORTRAN. How-

ever, it is rather cumbersome to automatically generate whole programs that way. For such purposes one better uses a specialized package like GEN-TRAN [49]. Another important problem is the optimization of the generated code which is usually necessary. In REDUCE the package SCOPE [128] was developed for this purpose. Its main strategy is to detect common subexpressions of large expressions.

MuPAD provides a very efficient form of interfacing, namely *dynamical modules* [121]. These are modules developed in a language like C or C++ that can be linked dynamically at run-time to the computer algebra system. Compared with approaches based on interprocess communication this leads to much less overhead. And as the module has direct access to the internal data of the *MuPAD* session, much less data must be communicated. The procedures implemented in a dynamical module can be called within *MuPAD* like any other function. Provided a convenient interface exists this allows in principle to work interactively with a numerical library.

Computer algebra systems can also help to *select* an appropriate method from a numerical library. Modern libraries have reached such a level of sophistication that for many users it is increasingly difficult to fully exploit their potential. They provide many different routines for the same task and the working of these routines can be further tuned by many input parameters whose meaning remains a secret for non-experts. A computer algebra system can analyze the given differential equation (e.g. estimate its stiffness) and then choose an appropriate method and determine reasonable values for its parameters. An example for this approach is the AXIOM package ANNA developed by Dupée [33].

Goldman *et al.* [51] go considerably further in their application of computer algebra by using it as a software engineering tool. They automatically *generate* the full FORTRAN code for numerically solving the Navier-Stokes equations. Their argument is that such programs are so long and complicated that their maintenance and adaption (new boundary conditions, different discretizations etc) is rather difficult and error-prone. They use instead a number of input files that contain all the relevant information about the problem in a format that is comparatively easy to read and let the computer algebra system then generate the complete code.

One can also use computer algebra to *derive* numerical schemes. The Butcher theory of Runge-Kutta methods is here a typical example. For higher order methods the order conditions become rather large and complicated. Computer algebra packages have been developed that derive and solve them (using Gröbner bases) [54, 120]. For partial differential equations the construction of higher-order discretizations or finite elements can be rather involved and is sometimes only feasible with the help of a computer algebra system [85].

At a more theoretical level computer algebra is used for the *analysis* of numerical methods. It may, for instance, assist in proving the stability of

finite difference schemes for partial differential equations [44]. Another example is the derivation of so-called modified equations [4], i.e. of differential equations whose solution is much closer to the numerically computed one than the solution of the original equation.

Another topic where computer algebra plays a certain role in numerical analysis are *differential algebraic equations*. The *index* of such a system comprising differential and algebraic equations measures in a certain sense, how far it is away from a pure differential equation [18]. This gives an indication of the difficulties one must expect in a numerical integration. The determination of the index is essentially equivalent to the completion procedures described in Sect. 5 [71, 95, 112], as it can be defined as the number of steps needed for the completion. However, in practice numerical analysts often prefer the use of *automatic differentiation* to computer algebra [19].

None of the applications described in this section represents really what one would call a *hybrid* algorithm combining symbolic and numerical elements, i.e. where computer algebra is an integral part of the solution process and not only used to determine either the problem or the numerical method for its solution.⁵ We are not aware of any such algorithm for differential equations, although some ansätze based on symmetry theory have been developed. Dorotnitsyn [32] showed how one may construct finite difference schemes inheriting the symmetries of a differential equation. Such schemes should very well preserve the associated conservation laws, but so far no numerical tests have been published.

In contrast, for solving algebraic equations several hybrid algorithms have already been designed. One of them deals with nonlinear systems possessing symmetries [47], as they e.g. arise in equivariant systems (see Sect. 8). In the symbolic part it uses the linear representation theory of finite groups to transform the problem into an optimal form for the numerical part. This includes for example a block diagonalization of the Jacobian. The numerical algorithm is complicated due to the underlying group theory. Gatermann [45] showed how the numerical computations can be automated by first computing the necessary group theoretic data which is summarized in a bifurcation graph.

10 Conclusions and Outlook

The application of computer algebra to differential equations is a vast field. We could only briefly discuss some of the main research directions and had to omit many others. For example, one can extend the idea of transforming differential equations far beyond simple heuristics and is then lead to the *equivalence problem of Cartan* [65, 88]. Within the algebraic approaches we

⁵One may, however, consider the work of Kleczka *et al.* [66] on the Poincaré map (see Sect. 8) as such a hybrid algorithm. See also [67].

ignored the theory of \mathcal{D} -modules [76] which is important in control theory.

The fields we have touched on are in rather different states. Some of them like symmetry theory are meanwhile fairly mature with the fundamentals well understood and they provide standard techniques for tackling differential equations implemented in many computer algebra systems. Others are still in an early stage of their development and essential questions are open. Such fields are usually known only to some experts and only prototypical implementations of algorithms exist.

One common feature shared by most of the fields is the complexity of the algorithms. If we take the various completion methods as example, it is obvious from their close relation to Gröbner bases that their (worst case) complexity is at least as bad as that of Buchberger's algorithm, i.e. double exponential. Although Gröbner bases solve *in principle* many problems in commutative algebra, it is well-known that one often fails to get a basis in reasonable time. One possible way out is the stronger use of heuristics and techniques from Artificial Intelligence, although this is an unpleasant thought for many pure mathematicians.

Some readers might be surprised that we discussed the combination of symbolic and numerical computations at such length. But in the future this topic will be among the most important ones — at least for applications. In the form of simple interfaces it happens already now in many places and hopefully we can soon add powerful hybrid methods. For most *users* of computer algebra systems (this is a very different community than the participants of computer algebra conferences!) such possibilities are of much greater importance than the fancy algorithms developed by theorists.

Despite all the successes of Lie symmetries, differential Galois theory etc. one must clearly see that these theories are of hardly any value for many of the problems an engineer for example typically faces. A popular benchmark problem in differential algebraic equations models with five links a car wheel suspension [59]. Its equations of motion must be generated by computer and consist of about 7000 lines of FORTRAN code. It appears hardly realistic to solve such a system by analytic techniques.

This does not imply that there is no point in studying symbolic methods. Toy models that can be solved analytically are very important for obtaining a deeper understanding of underlying structures. One may hope that such understanding may lead to more efficient numerical algorithms for such large problems. And again we want to stress that the application of computer algebra to differential equations is not restricted to solving them!

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